betaPrimePrecipitate tutorial

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1 Parameters

This section describes the model parameters that the user can access through the parameters.h file in each application folder. These can be divided between model-specific parameters and those generic to all models, and there are three categories of model-specific parameters: constant coefficients and tensors, bulk free energy functions, and residuals. In the context of their model, constant coefficients are self-explanatory. Bulk free energy functions and their derivatives are incorporated as functions of the phase field variables, usually instantiated in the code as n or c. All of the examples and defaults use polynomial free energies, and their derivatives are found symbolically. Non-polynomial functions could be implemented, but are not supported at this time. Residuals are used directly in evolution. Changing them is not recommended unless the user is comfortable with deal.ii and the PRISMS framework.

1.1 Generic Parameters

- problemDIM
 - Dimension of the problem (e.g. 1D, 2D, 3D)
- spanX

Length of system in x-direction

spanY

Length of system in y-direction. Not used if problemDIM < 2

• spanZ

Length of system in z-direction. Not used if problemDIM < 3

• refineFactor

Defines the refinement of the mesh. There are $2^{\text{refineFactor}}$ elements in each direction in this implementation, and $\left(2^{\text{refineFactor}}\right)^{\text{problemDIM}}$ elements in total.

• finiteElementDegree

The order of interpolation of the finite element space. In this case, the order of the Lagrange elements to be used.

• dt

The simulation timestep.

• numIncrements

The number of simulation iterations. Final time is then dt.numIncrements.

• writeOutput

Whether we are writing any output. Takes a boolean argument, e.g. true.

• skipOutputSteps

Output will be written every skipOutputSteps iterations. If writeOutput true, the initial conditions will always be written.

1.2 coupledCahnHilliardAllenCahn Parameters

This application implements the following set of equations for a conserved order parameter, c, and one or more non-conserved (structural) order parameters η :

$$\frac{\partial \eta}{\partial t} = -M_{\eta} \left(\frac{\partial f(\eta, c)}{\partial \eta} - K_{ij} \nabla_i \nabla_j \eta \right) \tag{1}$$

$$\frac{\partial c}{\partial t} = \nabla \cdot M_c \nabla \left(\frac{\partial f(\eta, c)}{\partial c} \right) \tag{2}$$

Since there is no gradient energy term in c, this might also be called a coupled Allen-Cahn/diffusion system. From a finite element perspective, Eq. 5 is more tractable as

$$\frac{\partial c}{\partial t} = \nabla \cdot M_c \left(\frac{\partial^2 f(\eta, c)}{\partial c^2} \nabla c + \frac{\partial^2 f(\eta, c)}{\partial \eta \partial c} \nabla \eta \right)$$
(3)

• numStructuralOrderParameters

Multiple structural/non-conserved order parameters can be used in this model (implemented?????)

• Mc

 M_c , mobility of the conserved order parameter c.

• MnVals

 M_{η} , mobility of the non-conserved order parameter η . This is an array with numStructuralOrderParameters elements, each corresponding to an order parameter.

• KnVals

 K_{η} , gradient energy coefficients K of the non-conserved order parameter η . Again, an array with numStructuralOrderParameters elements, each corresponding to an order parameter.

The free energy $f(\eta, c)$ is composed of free energies $f_{\alpha}(c)$ and $f_{\beta}(c)$ for the phases corresponding to $\eta = 1$ and $\eta = 0$, and an interpolating function $h(\eta)$. The default case uses free energies for a Mg-Gd system:

$$f_{\alpha}(c) = -1.6704 - 4.776c + 5.1622c^2 - 2.7375c^3 + 1.3687c^4 \tag{4}$$

$$f_{\beta}(c) = -1.5924 - 5.9746c + 5.0c^2 \tag{5}$$

These free energies should be in the form of a polynomial for this application. An interpolation function for a single order parameter should satisfy h(0) = 1, h'(0) = 0, h(1) = 1, and h'(1) = 0, for example:

$$h(\eta) = \eta^3 (10 - 15\eta + 6\eta^2) \tag{6}$$

The full bulk free energy (for n structural order parameters) is then:

$$f(\eta, c) = f_{\alpha}(c) \sum_{i=1}^{n} (1 - h(\eta_i)) + f_{\beta}(c) \sum_{i=1}^{n} h(\eta_i)$$
 (7)

Equations (7-9) and their derivatives are included in parameters.h as follows:

$$\begin{split} \text{faV} &= f_{\alpha}(c) \\ \text{facV} &= f_{\alpha}'(c) \\ \text{faccV} &= f_{\alpha}''(c) \\ \text{fbV} &= f_{\beta}(c) \\ \text{fbcV} &= f_{\beta}'(c) \\ \text{fbccV} &= f_{\beta}''(c) \\ \text{hV} &= h(\eta) \\ \text{hnV} &= h'(\eta) \end{split}$$

These free energy variables are incorporated into the residuals in the weak form of equations 4 and 6.

$$\begin{split} \texttt{rcxV} &= \left((1 - h(\eta)) f_\alpha''(c) + h(\eta) f_\beta''(c) \right) \nabla c + h'(\eta) (f_\beta'(c) + f_\alpha'(c)) \nabla \eta \\ \\ \texttt{rnV} &= (f_\beta(c) - f_\alpha(c)) h'(\eta) \\ \\ \texttt{rnxV} &= \nabla \eta \end{split}$$

1.3 betaPrimePrecipitate Parameters

The betaPrimePrecipitate application combines several features of other applications together to model evolution of coherent precipitates. In practical terms, this application extends coupledCahnHilliardAllenCahn by incorporating the strain energy of the transformation into the evolution of the structural order parameter. Following the Landau theory of phase transformations, a stress-free strain tensor (SFST) is defined for each orientation variant (structural order parameter) to be used in the simulation. The total stress-free strain tensor at a point \mathbf{r} , $\epsilon_{ij}^0(\mathbf{r})$, is the sum of each order parameter's SFST times it's interpolation function:

$$\epsilon_{ij}^{0}(\mathbf{r}) = \sum_{p=1}^{n} \epsilon_{ij}^{0}(p) h(\eta_{i}(\mathbf{r}))$$
(8)

Incorporation of this into the evolution equations is left to the corresponding model documentation.

- Mc, MnVals, and KnOTensor are inherited as is from coupledCahnHilliardAllenCahn.
- Ev, nuV, muV, lambdaV, and CijklV are as in mechanics

• sf0StrainV

This the stress-free strain tensor (a 9-element vector or 3×3 array) associated with order parameter 0. One SFST is defined for each structural order parameter.

$\bullet \ \mathtt{skipElasticitySteps}$

The elasticity problem is only re-solved after this many time steps of phase field evolution. To re-solve at every phase field iteration, set this equal to 1.

- faV, facV, faccV, fbV, fbcV, and fbccV are bulk free energy polynomials (and their derivatives) as in coupledCahnHilliardAllenCahn.
- h0V and h0nV are the interpolation function, $h(\eta)$ and its derivative $h'(\eta)$, as in coupledCahnHilliardAllenCahn, for order parameter 0. There is an interpolation function defined for each structural order parameter.

• Residuals

Residuals are defined as in coupledCahnHilliardAllenCahn, extended for multiple structural order parameters. More information about the derivation of these can be found in the accompanying Model documentation.

2 Initial Conditions

Initial conditions for the phase field parameters are generated in the functions InitialConditionN or InitialConditionC located in the main.cc file in the application folder. Two types of initial conditions have been implemented so far: a field of random noise around some mean value, and analytical functions of position. Random noise is the default for at least one variable in all of the phase field codes. Here, as an example, c is initialized to random values in the range 0.019-0.021:

```
template <int dim>
double InitialConditionC<dim>::value (const Point<dim> &p, const unsigned int
    /* component */) const
{
        //set result equal to the concentration initial condition
        return 0.02 + 1.0e-3*(2*(0.5 - (double)(std::rand() % 100 )/100.0));
}
```

Analytical functions of position can be generated from the members of the class p, e.g. p.distance or the x and y coordinates, p[0] and p[1]. In the default cases, the hyperbolic tangent of the signed distance from a circle or sphere is used. In the following example, η will be initialized into a sphere with radius=spanX/8, where $\eta=0.5$ at the boundary, $\eta\sim 1$ in the interior, and $\eta\sim 0$ outside.

```
template <int dim>
double InitialConditionN<dim>::value (const Point<dim> &p, const unsigned int
    /* component */) const
{
    //set result equal to the structural order paramter initial condition
    double dx=spanX/std::pow(2.0,refineFactor);
    double r=0.0;
    r=p.distance(Point<dim>(spanX/2.0,spanY/2.0,spanZ/2.0));
    return 0.5*(1.0-std::tanh((r-spanX/8.0)/(3*dx)));
}
```

This example requires that problemDIM=3. In the actual applications (e.g. coupledCahnHilliardAllenCahn), a conditional tests problemDIM, and there are analogous cases for each dimension.

3 Boundary Conditions

Boundary conditions are no-flux for the phase field evolution equations. Implementation of alternate boundary conditions is an objective for future releases.